
Temporal variability of dissolved trace metals at the DYFAMED time-series station, Northwestern Mediterranean

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Abstract :

We present here results of an 18-month survey (July 2007–March 2009) of a suite of selected trace metals (TM: Co, Ni, Cu, Pb) in a 2350 m-deep offshore water column in the Ligurian Sea (Northwestern Mediterranean Sea). This low-nutrient low-chlorophyll region is characterised by a long stratification period (May–November) during which surface waters are depleted of macronutrients. Trace metals exhibit a range of biogeochemical behaviours from surface-enriched (scavenged-type) to surface-depleted (nutrient-like) with Co and Ni as representative cases. Cobalt (28–172 pM) distributions are governed by external inputs of aeolian dust deposition and removal by adsorption onto particles in surface, intermediate and deep waters as well. Nickel (3.57–5.52 nM) distributions are governed by internal biogeochemical cycles, together with physical mixing and circulation patterns. Nickel is primarily removed from surface waters with biogenic particles and then remineralised at depth. Copper (1.39–2.89 nM) distributions illustrate a mixture of the two typical behaviours mentioned above. Distributions of typically anthropogenic and particle-reactive Pb (82–235 pM) are in agreement with a Mediterranean flow source of Pb for the adjacent North Atlantic Ocean. The mechanisms controlling the biogeochemical cycling of TMs, such as atmospheric inputs, physical forcing, and interactions with primary production, are discussed according to the TM physico-chemical properties and biological importance.

Highlights

► We sort out the influence of these different parameters in the water column in a low-nutrient low-chlorophyll region, the Ligurian sea (Western Mediterranean); ► The intensity of winter mixing determines the homogenization of TM concentrations in the water column; ► TM exhibit a range of biogeochemical behaviours from surface-enriched (scavenged-type) to surface-depleted (nutrient-like) with Co and Ni as representative cases; ► Atmospheric input is a chief parameter as is determine Pb, Cu and Co distributions, and preclude any limitation of primary production by TM; ► Anthropogenic Pb decreased by up to 60% and 20% in surface and deep waters the last 20 years, but water column is still Pb contaminated, in such a way that the Western Mediterranean deep-water outflow is the source of Pb contamination observed at 1000 m in the adjacent Eastern North Atlantic Ocean.

Keywords : Ligurian Sea, Trace metals, Seasonal variation, Mediterranean, Open water

1. Introduction

Several trace metals (TMs), such as Fe, Co, Ni, Cu, are oceanic micronutrients that are essential to marine life (e.g., Morel and Price, 2003), while others, such as Hg or Pb, are contaminants, and evidence of the large-scale human footprint on the ocean (e.g., Hatje et al., 2018). The distributions and residence times of both TM groups in the open ocean are regulated by external inputs (aeolian deposition, riverine and submarine groundwater inputs, hydrothermal activity, pore water release) and the rates of geochemical and biologically mediated (“biological pump”) removal processes (e.g., Geibert, 2018; Lohan and Tagliabue, 2018; Wangersky, 1986). The intensity of the biological pump is, in turn, dependent on physical processes that govern primary productivity at the surface. In conjunction with the variations of TM inputs, this latter process may generate a seasonal variability of TM concentration profiles, the observations of which are sparse due to the difficulty in producing offshore TM time series (Bown et al., 2017; Connelly et al., 2006; Saito and Moffett, 2002; Tappin et al., 1993).

Here, we present the first time-series (over the duration of 18 months) of TM distribution

over the water column in the central Ligurian Sea, in the Western Mediterranean, that allows characterisation of biogeochemical behaviours of four typical TMs (Co, Ni, Cu and Pb) in an oceanic environment where the dynamics of water circulation drive a high biogeochemical seasonality. Due to the occurrence of strong physical forcing such as horizontal and vertical advection on open waters (Béthoux, 1989), and, even more, the prominent role of the atmospheric deposition (Migon et al., 2002), the Mediterranean Sea is a suitable oceanic environment for studying seasonal dynamics of TMs.

2. Site description in the Mediterranean context

The Mediterranean thermohaline circulation is characterised by the inflow of Atlantic water at the surface and exiting below Mediterranean Overflow Water (MOW) at Gibraltar. An important consequence of this circulation is the short residence time of Mediterranean waters (50 to 100 years on average; Millot and Taupier-Letage, 2005), compared to that of other ocean basins (200 to 1000 years; Durrieu de Madron et al., 2011). Owing to the peculiar characteristics of the basin (i.e. shallowness of the Sicily Strait with 316 m maximum depth, evaporation and rain budgets, dense and bottom water formation), the mean water residence time of the Western Mediterranean Sea is even shorter, amounting to 22 ± 4 years (Roether et al., 2013). The process of dense water formation ventilates deep waters during winter (Testor et al., 2018; Waldman et al., 2018). In addition, the Western Mediterranean Sea has relatively small dimensions (e.g., the Ligurian Sea: $5.3 \cdot 10^{10} \text{ m}^2$), and it is subjected to significant anthropogenic pressure and pulsed natural Saharan dust events (Moulin et al., 1997). In particular, atmospheric deposition becomes the major source of input of TM and nutrients to the surface photic layer during the stratification period (Bartoli et al., 2005; Davis and Buat-Ménard, 1990). This results in significant fluxes of various elements and compounds, including trace

metals (TMs). To a large extent, the atmospheric TM inputs govern their offshore biogeochemical cycling (Guerzoni et al., 1999; Migon et al., 2002).

The DYFAMED (DYNAMIQUE des Flux Atmosphériques en MEDiterranée; Coppola et al., 2016) sampling site is located in the central zone of the Ligurian Sea, ~ 50 km off the continental French coast, 2350 m deep, 43° 25'N, 7°52' E (Fig. 1). This site is surrounded by the permanent geostrophic Ligurian frontal jet flow that results from the cyclonic circulation of the North Current, formed in the Ligurian Sea by joining the currents flowing northward along the coasts of Corsica (Millot, 1999). The North Current creates a band (~30 km width, > 250 m depth) that presumably shelters the sampling area from riverine and coastal lateral inputs by a strong horizontal density gradient (Niewiadomska et al., 2008). Acoustic Doppler Current Profiler (ADCP) measurements have shown that the DYFAMED station is isolated from the Northern Current (Andersen and Prieur, 2000), even if infrequent intrusions of waters may episodically occur in winter (Millot, 1999). As a result, the site is believed minimally affected by lateral inputs, and it is therefore commonly used as a reference site for the impact of atmospheric inputs to an oligotrophic open-ocean water column (Martín et al., 2009; Migon et al., 2002; Sarthou and Jeandel, 2004). The DYFAMED site is regarded as a one-dimensional station where simple hydrological mechanisms prevail and where the ecosystem is quite well understood (Avril, 2002). Owing to these features and its accessibility (only 50 km from Nice), the site has been chosen as a time-series observation station and has been monitored monthly since January 1991 (Marty, 2002). The site has been extensively studied for ~30 years, e.g., DYFAMED and MEDFLUX programmes; see special issues *Deep-Sea Research II* 49, 11 (2002) and 56, 18 (2009), respectively. The DYFAMED station is now part of the Mediterranean Ocean Observation Site for the Environment (MOOSE; <http://www.moose-network.fr>).

Three major water masses can be distinguished in the Ligurian Sea (Marty and Chiavérini, 2010; Millot, 1999): the surface water (0–200 m-depth), strongly affected by local climatic conditions (Schröder et al., 2006), the subjacent layer (approximately between 200 and 600 m depth at the DYFAMED site), characterised by maximum temperature and salinity due to the intrusion of Levantine Intermediate Water (LIW) (Marty and Chiavérini, 2010; Millot, 1999), and the deep layer, namely the Western Mediterranean Deep Water (WMDW), characterised by relatively low salinity and low temperature. Above all, the Ligurian Sea is subject to pronounced seasonal variations in hydrology and biology (Marty et al., 2002), with a long stratification period (May–November) during which nutrients are depleted in the photic layer. During the stratification period, atmospheric deposition becomes the major source of inputs of TMs and nutrients to the surface photic layer (Bartoli et al., 2005; Migon et al., 2002). The stratification is then disrupted in late autumn, and, in winter, due to a strong physical forcing (action of cold and dry winds, combined with low temperatures), surface waters are cooled, and their evaporation is enhanced, increasing their salinity. This process regularly initiates the formation of dense water. The subsequent vertical mixing is likely to bring amounts of nutrients from depths to surface layers, which, thus, enable early spring phytoplankton blooms (D’Ortenzio and Ribera d’Alcalà, 2009; Heimbürger et al., 2013; Migon et al., 2002). Dense water formation involves the oxygenation of deep layers. Part of this oxygen is consumed by the remineralisation of organic matter (OM) produced and settled from the photic zone.

Seasonal dynamics of phytoplankton at DYFAMED have been characterised using pigment analysis (Marty et al., 2002). The 0–200 m-depth integrated total chlorophyll a ([TChl-a]) is highly variable throughout the year and reaches the highest values during the spring bloom (up to 230 mg m⁻²) and, evidently, the lowest values during the period of

the stratification period (down to 12 mg m^{-2}). Dissolved OM and TMs generally accumulate within the photic layer during summer stratification and are then exported to deeper layers (Avril 2002; Copin-Montégut and Avril, 1993; Heimbürger et al., 2014; Migon et al., 2002).

3. Material and Methods

3.1. Sampling

In order to catch, during our sampling period, one seasonal cycle, observations over a period of at least one year and a half were required (Granger, 1978). Thus, monthly vertical profiles of TM and ancillary parameters were established over the duration of 18 months at the DYFAMED station, between June 2007 and March 2009, except in December 2008 and February 2009, due to bad weather conditions. On each cruise aboard the RV *Téthys II*, seawater was generally sampled at 12 depths from surface to 2200 m (150 m above the seafloor), including the main features such as the surface mixed layer, TChl-a maximum, LNV, and WMDW. Hydrographic parameters were obtained using a CTD profiler (Sea-bird SBE 911plus) with additional sensors (dissolved oxygen and Aq-tracker MKIII Chelsea fluorometer). Water samples were taken using trace-clean 5-L Teflon-coated Niskin X bottles (General Oceanics 1010X) fixed on a carousel water sampler (Sea-bird SBE32) that was coated with a metal-free epoxy-based paint, to allow trace-clean sampling. Prior to sampling, Niskin X bottles were carefully cleaned in a clean room (Class 100) with diluted nitric acid washes (Suprapur HNO_3 , Merck) and deionised water rinses (Milli-Q, Millipore, resistivity $18 \text{ M}\Omega \text{ cm}$). Unfiltered samples were collected along the entire water column, while filtered sub-samples were obtained only from the upper 50 m, where most particles and phytoplankton reside (Niewadomska et al., 2008). In absence of any measurement of

particulate TM concentration in Mediterranean waters, we extrapolated data from other places in the world ocean (Nakatsuka et al., 2007). We calculated that, at depths > 50 m, the particulate fraction of TMs is < 3 % and may thus be considered negligible.

Filtrations were performed under high-purity nitrogen pressure (1-1.2 bar), in line with the Niskin bottles through filter cartridges (Sartobran 300, 0.2 μm). Sub-samples for TM measurements were immediately drawn into acid-cleaned 500 mL polyethylene (PE) bottles following the GEOTRACES trace-clean sample handling protocols (compatible with the www.geotraces.org/images/Cookbook.pdf, version 3, 2017). They were then acidified with HNO_3 (Suprapur, Merck) to pH 1. All PE bottles were hermetically sealed, double-wrapped in polyethylene bags, and kept at +4°C until analysis.

3.2. Chemical analyses

Filtered sea water samples were extracted in the clean laboratory of Centre Ifremer Atlantique following Danielsson et al. (1982). Briefly, the method can be described as chelation of the dissolved TMs with a mix of ammonium pyrrolidine dithiocarbamate (APDC) and diethylammonium N,N-diethylcarbamate (DDDC), extraction of the chelates with Freon, and back-extraction with diluted HNO_3 . The obtained solutions were then analysed using an inductively coupled plasma-mass spectrometer (ICP-MS, Thermo Electron Corporation, Element X Series[®]). The detailed procedure for the extraction is described elsewhere (Chiffoleau et al., 2004). Precision and accuracy were tested using two Certified Reference Materials (CRM) from the National Research Council of Canada, namely coastal waters (CASS-1 to 3) and open ocean water (NASS-2). Precisions, calculated as the variation coefficient (i.e. confidence interval/mean) of 6 replicate analyses of CRMs, varied between 2 % and 9 % (Table S1, Supplementary Information, SI). The detection limits were estimated 3 times the standard deviation of

the 6 blank replicates obtained from an analytical run. They were 8, 20, 60, and 2 pM for Co, Ni, Cu, and Pb, respectively, which is low compared to the range of determined concentrations (Table S2), and generally to typical oceanic concentrations (Schlitzer et al. 2018). Accuracies of analytical results were tested upon the results of CRM analyses; they were always within the target range for certified values for the four TMs analysed. Sampling quality was tested upon oceanographic consistency of the TM distributions. No outliers were noticed for Co, Ni and Cu. Only a few samples, taken at the beginning of the sampling period, exhibited very high Pb concentrations compared to the adjacent samples. This excess decreased as the sampling programme proceeded. As they were without any oceanographic consistency, we removed them from the data set (Table S2, SI). Phosphate (PO_4^{3-}) and silicate ($\text{Si}(\text{OH})_4$) concentrations were determined using a standard auto-analyser according to the protocol developed by Aminot and K erouel (2007), with detection limits of $0.02 \mu\text{M}$ for both phosphate and silicate, and precisions better than 6%. Pigment determination was performed using high-performance liquid chromatography (HPLC) techniques described in Ras et al. (2008), $\int\text{TChl-a}$ being the sum of chlorophyll-a + divinyl-chlorophyll-a concentrations integrated within the upper 200 m of the water column. The density gradient between the surface (10 m) and the base of the mixed layer (ML) estimated to determine the mixed layer depth (MLD) was 0.03 kg m^{-3} (de Boyer Mont egut et al., 2004; D'Ortenzio et al., 2005). For data interpretation and associated statistics, we separated samples collected above 100 m depth, and samples collected below 100 m depth, in order to compare TM distributions in two layers that have contrasted concentration profiles. This approach is justified because the MLD never extended beyond 100 m during the present sampling period (Pasqueron de Fommervault et al., 2015).

3.3. Statistical analysis

Parametric (Student t-test, Pearson correlation) and non-parametric (Wilcoxon, Mann-Whitney) statistical tests were applied to variables after testing for the normality of distributions (Shapiro-Wilk and Fisher tests). The statistical computations were performed with XLSTAT software from Addinsoft (<https://www.xlstat.com/>). The frequency distributions of TM concentrations showed some departure from normal-distributions (Fig. S1 and Table S3, SI). Thus, in order to describe central tendency measurements, we calculated geometric means and median, in addition to arithmetic means and standard deviations (Table 1).

4. Results and Discussion

4.1. Hydrological and biological seasonal patterns

Salinity varied between 38.02 and 38.62, within the range usually found at this station (Marty et al., 2002). Vertical salinity profiles always followed the well-known pattern for this site, with lowest values in the ML, maximum values between 200 and 600 m (LIW) and moderate values below 600 m (WMDW). Temperature ranged from 12.9 to 24.6°C, with highest values in the shallow ML (~10 m) at the end of summer. Vertical temperature profiles converged below 1000 m-depth to approximately 12.9°C, while the overlying LIW was slightly warmer with approximately 13.4°C. Seasonal patterns of MLD during the sampling period are shown in Fig. 2 (see also Pasqueron de Fommervault et al., 2015). The water column remained well stratified most of the time, and only slight winter mixing occurred. In contrast to usual MLD values of 100 - 200 m (Heimbürger et al., 2013; Marty et al., 2002) or more (> 2000 m-depth in 2006; Marty and Chiavérini, 2010) during winter mixing, the MLD never exceeded 70 m-depth during our sampling period (Fig. 2). Among other physical forcing factors, MLD is viewed to have the strongest impact on phytoplankton dynamics (Lavigne et al., 2013).

The relatively shallow MLD observed during our sampling period may not be deep enough to supply sufficient macronutrients and, therefore, may be responsible for the low primary productivity resulting in low values of \int TChl-a. However, TChl-a varied intensely over the sampling period (Fig. 2); its distribution reveals spring and autumn phytoplankton blooms within the upper 50 m (Fig. 2). Values of \int TChl-a were in the range of those usually found (12 - 230 mg m⁻²) at this site (Marty et al., 2002), ranging between 16 and 170 mg m⁻².

An outstanding feature is the high \int TChl-a value in October 2007 (77 mg m⁻²). Nutrients from subsurface layers were likely raised to the surface layer by the permanent cyclonic circulation of the area (D'Ortenzio et al., 2014). Due to the relative shallowness of nutriclines in this region, wind events can inject some nutrients to the depleted surface layer, hence an increase of chlorophyll concentration. Furthermore, exceptionally high \int TChl-a values were associated to the period when winter convection takes usually place (January-February 2008, 102 and 123 mg m⁻², respectively; Fig. 2), in spite of a very shallow MLD measured in winter 2008 (Coppola et al., 2018). Before the use of autonomous platforms capable of acquiring high-frequency data (i.e. the time of our sampling), MLD measurements were made during monthly on-site campaigns scheduled a year in advance, involving recurring uncertainty in the measurement of a process of relatively short duration. Hence the possibility of incorrect estimates of the MLD, likely to yield errors in the concentration of nutrients brought to surface waters by the winter convection process. Such incorrectly estimated (underestimated) MLD can lead to a mismatch between the calculated nutrient supply, the amount of Chl-a that can be sustained by such a supply, and the actual observed Chl-a concentration. After the spring bloom in April 2008 (\int TChl-a = 170 mg m⁻²), \int TChl-a remained greater than 40 mg m⁻² until September 2008.

Figures 3a and b present the vertical PO_4^{3-} and Si(OH)_4 concentration distributions, respectively, over the entire water column for the whole sampling period. The general shape of these profiles was typical of the distribution usually observed at the DYFAMED site (Marty et al., 2002), with highest concentrations between 500 and 1000 m-depth, especially between March and July 2008, and lowest concentrations in the surface layer. Deepening of the ML, in March 2008 and January 2009, down to 41 and 69 m, respectively (Fig. 2), lead to partial homogenisation, after which slightly higher macronutrient concentrations were present in the ML, in March 2008 and 2009, that is 0.08 to 0.15 μM for PO_4^{3-} and 1.4 to 3.7 μM for Si(OH)_4 (Fig. 3).

In summary, our sampling period from mid-2007 to mid-2009 displayed (i) a highly stratified water column with only little dense water formation in winter (based on MLD values measured at DYFAMED), and (ii) phytoplankton blooms in spring and late summer-early autumn (especially autumn 2007).

4.2. Trace metal distributions

The most striking characteristic of the TM distribution in the Ligurian Sea over the 18-month sampling period is the high concentration of Pb. On the contrary, when comparing the present data with those of the Schlitzer, eGEOTRACES – Electronic Atlas (Geotraces Electronic Atlas, 2019), it appears that Co, Cu and Ni concentrations were within the ranges usually found in open oceanic waters of the Mediterranean Sea (Schlitzer et al. 2018). Historical literature data are recorded in Table S4 (SI) for comparison. While most TMs changed little over time, Pb concentration decreased continuously during the last 20 years (see discussion below). Major variations of the present TM time series occurred in the upper 100 m (Fig. 4), where physical, chemical and biological forcing is the most intense. Below 100 m, most TM concentration profiles converged rapidly toward a relative uniformity, indicating a rapid mixing time

or a common export mechanism with little temporal variability. Basic processes occurring in the water column can be assessed by running a simple vertical two-box model, separating the euphotic layer (< 100 m) from the deep reservoir (> 100 m) at DYFAMED. The four TMs studied have significantly different mean concentrations ($p < 0.01$) between the euphotic (< 100m) and deeper waters (> 100m) (see statistical tests in Table 1). On the one hand, mean concentrations were higher in the deep waters than in the euphotic layer for Ni and Cu, confirming their “nutrient-like” behaviour. On the other hand, mean concentrations were higher in the euphotic layer than in deep waters for Co and Pb, which confirms the “surface-enriched/scavenging” type of their vertical distribution, underlining the importance of atmospheric sources for these two TMs. We did not observe any increase in TM concentrations near the bottom that suggests negligible benthic TM source. This idea needs further consideration since our sampling did not include the bottom 150 m of the water column. This water layer may contain deep-sediment resuspension resulting from bottom-reaching winter convection, or deep cascading events occurring in Western Mediterranean Sea (Durrieu de Madron et al., 2017). Generally, there is no nepheloid bottom layer at DYFAMED, except for rare cases of very strong events of deep convection and cascading, e.g., in 1999 (Béthoux et al., 2002), or, even more, in 2006, when an exceptional episode of deep convection occurred (Martín et al., 2010). As already mentioned, the process of dense water formation was not very significant during the sampling period considered here, and the presence of a nepheloid layer is unlikely.

Cobalt concentrations varied from 22 to 172 pM, and were high in the upper 100 m (up to 172 pM at surface; Table 1) and decreased rapidly within the upper 200 m to fairly uniform concentrations of 41 ± 8 pM, a value very close to the 45 pM determined in the WMDW (Dulaquais et al., 2017). Such a distribution is typical of surface-

enriched/scavenged-type behaviour of TMs having a large atmospheric source. The winter surface concentrations are as high or higher than those measured in summer, i.e. around 100 pM (Fig. 4a). This might be due, in 2008, to the relatively late occurrence of the convection process (Coppola et al., 2018), i.e. in March. At the end of the spring bloom, dissolved Co concentration is minimal because of its consumption by plankton. As oligotrophy increases, Co consumption decreases and, at the same time, atmospheric deposition refuels surface waters. Thus, the stock of dissolved Co increases to presumably reach its maximum level before the convective export. In 2009, no significant mixing was observed at DYFAMED (Coppola et al., 2018). Consequently, our results did not show any pronounced seasonal difference in the general distribution pattern. Vertical Co profiles always conserved their surface-enriched shape, as observed in the entire Mediterranean Sea (Dulaquais et al., 2017). However, some seasonal pattern in the surface layer can be recognised. High Co surface concentrations (> 100 pM) were observed from October 2007 to March 2008, suggesting that above-average atmospheric loads of Co were deposited to surface waters and have led to the accumulation of dissolved Co between the end of the plankton bloom and the dense water formation episode, i.e. an oligotrophic period during which there is almost no export, due to the absence of biological activity to allow vertical transfer (e.g., Migon et al., 2002; Passow, 2004; Heimbürger et al., 2014). This ended when the process of dense water formation took place, allowing a rapid export of dissolved matter. As well, Co concentrations in the surface layer peaked up to 164 - 172 pM in January 2009 (Fig. 4a). Indeed, although no data on atmospheric deposition were available in 2009, very high Co concentrations were measured in the atmospheric aerosol (Cap Ferrat sampling station) in October 2008 (up to $11 \mu\text{mol m}^{-3}$ while the mean aerosol Co concentration was $1.8 \pm 1.3 \mu\text{mol m}^{-3}$ between January 2006 and December 2008; Heimbürger et al., 2010). Estimations of seawater-

labile Co inputs to the surface of the Ligurian Sea ranged between $0.08 - 6 \mu\text{mol m}^{-2} \text{d}^{-1}$ (Heimbürger et al., 2011). The increase of concentrations in the atmospheric aerosol observed in October 2008 suggests an increase of Co deposition fluxes as well. This atmospheric input occurred during the oligotrophic period, and its seawater-labile fraction was presumably transferred to depth only during the formation of dense water.

Furthermore, the statistically significant negative correlation between Co and salinity ($[\text{Co}](\text{pM}) = -187[\text{Salinity}](\text{psu}) + 7242$, $n = 204$, $R = 0.80$ for a salinity range comprised between 38.0 and 38.6 psu) suggests that the atmospheric Co source consists mainly of wet deposition, the influence of riverine discharge being very low in the Ligurian Sea, overall.

Lower concentrations ($< 100 \text{ pM}$) were observed from April to September 2008, in agreement with uptake by phytoplankton in spring. Cobalt is a bioactive TM of considerable importance in biogeochemical processes occurring in the global ocean (Morel, 2008), but also the scarcest of all known bioactive TMs (Saito et al., 2017). Owing to its surface-enriched pattern, the steep decrease of Co within the upper 200 m-depth should not be related to its consumption by PP, but may be attributed, at least partially, to adsorption onto sinking particles (Fisher et al., 1991) and further transfer to the marine sediment. In oligotrophic waters, the bioactive character of Co may be crucial, but it is very likely outweighed by the relative importance of atmospheric inputs in the Northwestern Mediterranean (Guerzoni et al., 1999; Marín et al., 2017). In spite of this important bioactive role, Co surface depletions are not usually seen unless Zn values are low, according to the hypothesis that Co may replace Zn in marine phytoplankton, when Zn becomes depleted (Morel, 2008; Sunda and Huntsman, 1995). Considering the relatively high values of Co and Zn in Northwestern Mediterranean waters (Migon, 2005), the occurrence of this substitution mechanism remains unlikely. When PP was

maximal (February-April 2008, March 2009), surface-enriched Co profiles were always conserved (Fig. 4a).

Cobalt is strongly bound by ligands, and the speciation study of Saito and Moffett (2001) suggests that a fraction of Co might not be detectable by many analytical methods without UV-oxidation procedure. This possible bias, not detectable by the analysis of CRM samples, might affect Co concentrations, but presumably not Co biogeochemical behaviour, as Co exhibits relatively high concentrations in surface, compared with oceanic regions, and its distribution is characterised by accumulation in surface waters. We infer that the distribution of Co in the upper water column is mainly controlled by steady inputs of atmospheric crustal material, while biological activity, even in mesotrophic conditions (spring bloom), is not capable of removing Co from surface waters to the point of surface depletion. In summary, the present data set suggests that Co does not co-limit PP in the Northwestern Mediterranean basin, because of high atmospheric inputs. Future increased large-scale use of Co for automotive batteries will likely further increase Co inputs to the Mediterranean Sea, and the Global Ocean as well (Zhang et al., 2016).

Nickel concentrations varied from 3.57 to 5.52 nM, with a mean of 4.54 ± 0.39 nM (Table 1). These concentrations are comparable to those previously published for the western Mediterranean (Table S4, SI, according to Morley et al., 1997 and Yoon et al., 1999), and vary only little with time and depth (Fig. 4b). The variability of the observed seawater values (8.6%) approaches the analytical precision (5%). However, the graphical illustration of the profile series suggests lower concentrations in the euphotic waters (Fig. 4b), which is attested by statistical tests ($p < 0.01$, Table 1). High PP in spring (2008 and 2009) lead to the lowest Ni surface concentrations (Fig. S3). Nickel displayed typical nutrient-like profiles throughout the sampling period. The low surface concentrations are

explained by biological uptake (Dupont et al., 2007), while higher deep-water concentrations are presumably due to the export flux of OM with a subsequent release during remineralisation. Indeed, Ni is significantly correlated ($p < 0.01$) with phosphate ($[\text{Ni}](\text{nM}) = 1.80 [\text{PO}_4^{3-}](\mu\text{M}) + 4.03$, $n = 180$, $R = 0.81$), and silicate ($[\text{Ni}](\text{nM}) = 0.09 [\text{Si}(\text{OH})_4](\mu\text{M}) + 4.03$, $n = 169$, $R = 0.77$), as previously pointed out by, e.g., Achterberg et al. (1993). The lowest surface Ni concentrations are presumably due to the consumption of the Ni stock during the spring bloom. In addition, it is likely that the uptake by PP is not balanced by atmospheric inputs, which are usually low in winter (Heimbürger et al., 2010). In all cases, Ni profiles were fairly uniform at depth, with no noticeable change in the different seasons, as already stated by Morley et al. (1997). In summary, the present data indicate that Ni distributions are largely controlled through uptake by microorganisms and remineralisation from settling particles, as already suggested by Cid et al. (2011).

Copper concentrations varied between 1.39 and 2.89 nM with a mean of 1.99 ± 0.28 nM (Table 1 and Fig. 4c). Values in the same range have been observed since the 1980s (Table S3, SI, according to Béchoux et al., 1990; Boyle et al., 1985; Morley et al., 1997). Although, on a yearly basis, Cu concentration were lower in the euphotic layer than deeper ($p < 0.01$, Table 1), surface-enriched Cu profiles (e.g., December 2007) alternated with surface-depleted ones (e.g., May 2008), and uniform profiles also occurred (e.g., March 2009). Surface depletion occurred when driving forces of vertical export (hydrology with dense water formation, or biology in productive conditions; Migon et al., 2002) were active, while surface accumulation occurred under summer/autumn conditions of oligotrophy with little uptake by PP. This scheme was not always clear, however. For example, in 2008, Cu vertical distributions shifted from surface-depleted profile (February) to surface-enriched profile (March), and then shifted again toward

surface-depleted profiles (April and May; Fig. 4c). Such a scenario may seem surprising, because the maximum deepening of the ML in 2008 was observed in March (Fig. 2), which suggests a hydrologically-driven transfer of matter to depth, thus no accumulation in surface. As Cu concentrations in the atmospheric aerosol were regularly decreasing between February and May 2008 (Heimbürger et al., 2010) the possibility of atmospheric loads capable of counterbalancing this transfer is unlikely. Therefore, we think the most plausible explanation is the inaccuracy of field data on the exact time of winter convection process as already pointed out by Heimbürger et al. (2013). More generally, Cu, in spite of its well-known biological role, rarely depicted nutrient-like profiles in the Northwestern Mediterranean, as observed in the global ocean, e.g., Pacific (Bruland, 1980; Nozaki, 1997). In the oligotrophic waters of the Northwestern Mediterranean, the bioactive character of Cu is probably outweighed by the relative importance of atmospheric inputs. However, biological uptake during spring bloom seems capable of consuming important Cu quantities, inducing a rapid shift toward surface-depleted profiles as observed as soon as in April 2008 (Fig. 4c). This strong dependence of Cu dynamics on biological activity (seasonality) highlights the importance of temporal observations. In particular for Cu, one has to take into account the current state of PP in the interpretation of a vertical Cu profile. For example, Boyle et al. (1985) and Seyler et al. (1989) found nutrient-like profiles for Cu just after the spring bloom. On the contrary, others reported a more or less homogeneous distribution, with, perhaps, a slight surface-enrichment (Béthoux et al., 1990; Ferrara and Seritti, 1989; Yoon et al., 1999). In summary, Cu profiles in surface waters seem close to the rectilinearity, presumably resulting from equilibrium between removal by plankton consumption and atmospherically-deposited loads. This would result in a shift between surface-depleted Cu profiles in periods of high biological activity and surface-enriched Cu in periods of

oligotrophy.

Lead concentrations ranged from 82 (below 100 m-depth) to 235 pM (above 100 m-depth) with a mean value of 117 ± 24 pM over the 0-2200 m-depth water column (Table 1). Lead concentrations depicted intense variation in the euphotic layer (Fig. 4d), but minor, although statistically significant, enrichment compared to the deep waters (Table 1), with a large variability of the concentrations in the water column (24%). Basically, Pb concentrations exhibit surface-enriched profiles. Compared to previous published data, Pb concentrations in the Western Mediterranean Sea have decreased over time in surface waters (0-100m, approximately) (Nicolas et al., 1994; Migon and Nicolas, 1998; Table S4, SI). Moreover, we found surface Pb concentrations similar to those found earlier in deep waters. This results from the decrease of external fluxes since mid-1980s, mainly atmospheric deposition (Migon et al., 1993; Migon et al., 2008). While atmospheric Pb concentrations in the Mediterranean troposphere decreased by 90% between 1987 and 2008 (Heimbürger et al., 2010), a smaller decrease was observed in seawater Pb concentrations. The Pb concentrations of the upper 100 m decreased by 60%, from 300-362 pM (Seyler et al., 1989; Marty and Nicolas, 1993) to 126 pM, and by 20% in waters below 100 m depth, from 130-145 pM (Seyler et al., 1989; Marty and Nicolas, 1993) to 108 pM respectively).

Using the mean concentration of deep Pb (108 pM, Table 1) as the typical Pb concentration in the MOW, this concentration is more than twice the concentration found in the North Atlantic at the same depth (< 40 pM according to Boyle et al., 2014).

Moreover, this is consistent with the recent results by Zurbrick et al. (2018), who found that Pb concentrations in the Atlantic off the Portugal coasts were the highest in subsurface Mediterranean waters, i.e., ~ 60 pM in the core of the MOW body. Likewise, increased dissolved Pb concentrations observed along GEOTRACES transects in the

Northeastern Atlantic were identified as MOW inputs (Rolison, 2016). Rusiecka et al. (2018) have described the spreading of MOW across Northeast Atlantic at intermediate depth.

4.3. Trace metal behaviours: sorting physical and biogeochemical processes

In summary, one can find at a single offshore site (DYFAMED), on a temporal scale, all the behaviours previously observed on a geographic scale. The four TMs (Ni, Cu, Co and Pb) exhibit a range of biogeochemical behaviours that can be characterised by two typical concentration profiles, surface-depleted (nutrient-like) and surface-enriched (scavenged-type). Nutrient-like metals, best exemplified here by Ni are primarily removed from surface waters by biogenic particles and then remineralised at depth. Internal biogeochemical cycles, together with physical mixing and circulation patterns, control the distributions of nutrient-like TMs. Scavenged-type metals, exemplified here by Co and Pb, are scavenged by particulate matter and continue to be removed onto particles in intermediate and deep waters as well as at the surface. External inputs, such as the deposition of aeolian dust, largely control the concentrations and distributions of scavenged-type TMs, which result from the equilibrium between atmospheric supply and removal by scavenging and sometimes uptake (e.g., Co). Overall, this equilibrium results in a surface enrichment, the magnitude of which varies over the season as the supply and removal respective intensities vary. Surface Cu distribution results from the balance between atmospheric supply and biological uptake. When supply outcompetes uptake, Cu accumulates in the surface layer. For Ni, the supply never leads to surface accumulation, but these two bioactive TMs both behave like nutrients rather than scavenged-type TMs, because deep concentrations of both are not depleted, which is the identifying feature of nutrient-like elements.

In the Mediterranean area, TMs with significant crustal/natural sources (Co, Ni, and, to a

lesser extent, Cu) are characterised by higher atmospheric inputs during summer, owing to pulsed Saharan dust emissions (Marticorena and Bergametti, 1996). On the contrary, atmospheric fluxes of anthropogenic TMs such as Pb are generally higher in winter (Duncan and Bey, 2004; Heimbürger et al., 2010; Migon et al., 2008). However, we could not observe any evident coupling of TM vertical distributions with seasonal variation of atmospheric TM inputs measured during the same period on the nearby coastal sampling site Cap Ferrat (Heimbürger et al., 2010). This might support the hypothesis that atmospheric deposition does not determine marine vertical export flux of metals, but accumulates in the marine surface layer, awaiting to be exported by hydrological or biological means (Heimbürger et al., 2014; Migon et al., 2002). Particle-reactive TMs are rapidly removed from the water column to the marine sediment. For example, Pb adsorbs onto particles (e.g., Bridgestock et al., 2018; Shen and Boyle, 1988) and co-precipitates on iron/manganese oxides (Tanguy et al., 2011; Waeles et al., 2007), hence its scavenging-type behaviour. Removal processes of such TMs are mainly governed by biological activity and their residence times (e.g., Cu and Pb, ~9 and 0.5 years; Nicolas et al., 1998) are generally lower than the WMDW residence time. Due to the rapid ventilation of Mediterranean waters, Pb concentrations evolved rapidly, almost concomitantly with atmospheric changes (Migon and Nicolas, 1998; Heimbürger et al., 2010). The signature of this evolution has been found in the MOW, which feed into the Atlantic Ocean (Noble et al., 2015; Zurbrick et al., 2018). However, Pb concentrations remain higher than in other oceanic environments of the northern hemisphere, suggesting the persistence of the anthropogenic emissions.

In summary, the 18-month time series presented in this paper allowed to exemplify the superimposed effects of water stratification, atmospheric input and biological pump on the vertical distribution of TMs in a particular open-sea system. The short residence

time of waters, the alternation of deep convection and water stratification, and the succession of different trophic states of the Ligurian Sea, make it a suitable environment to illustrate the entanglement of these processes. The present study allows to sort out the influence of biogeochemical and physical parameters on TM distribution in the water column of a low-nutrient low-chlorophyll region, the Ligurian Sea (Western Mediterranean). The Ligurian Sea is an epicontinental sea where deep waters are formed during winter convection. Our results showed that Co, Cu and Ni concentrations were within the ranges found in other open oceanic waters and were not limiting the primary production, whereas Pb levels characterised anthropogenic inputs, albeit its 20% concentration decrease recorded in deep waters in the last 20 years. Winter convection spreads Pb contamination in the deep waters of the entire Western Mediterranean, suggesting that the Mediterranean outflow is the source of Pb contamination, observed at around 1000 m depth, in the adjacent Eastern North Atlantic Ocean.

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Table caption

Table 1: Statistics for trace metal (TM) concentrations measured at the DYFAMED station (Ligurian Sea) from July 2007 to March 2009. Min.: minimum; Max.: maximum; N: number of determinations, SD: standard deviation. The last line of the table indicates the significance level of the difference in TM mean concentrations between surface-100 m and 100 m-bottom layers. T-tests were applied on normally distributed concentrations, i.e. Ni and Cu, whereas non-parametric Mann-Witney tests were performed for Co and Pb (see Fig. S1, in SI, for normality testing). Thus, when distinguishing photic (< 100 m) from deep layer (> 100 m), it appears that mean concentrations were systematically higher in the photic zone for Co, Cu and Pb.

	Co (nM)	Ni (nM)	Cu (nM)	Pb (pM)
<i>Full water column</i>				
Min. – Max. (N)	28 – 172 (204)	3.57 – 5.52 (200)	1.39 – 2.89 (187)	82 – 235 (130)
Arithmetic mean (SD)	61 (29)	4.54 (0.39)	1.97 (0.28)	117 (24)
Geometric mean (Median)	55 (51)	4.52 (4.58)	1.95 (1.93)	115 (112)
<i>Surface – 100 m</i>				
Min. – max. (N)	56 – 172 (88)	3.57 – 5.37 (90)	1.39 – 2.89 (81)	90 – 235 (63)
Arithmetic mean (SD)	88 (25)	4.26 (0.34)	2.07 (0.30)	126 (28)
Geometric mean (Median)	85 (81)	4.24 (4.22)	2.05 (2.03)	123 (120)
<i>100 m – bottom</i>				
Min. – max. (N)	28 – 70 (116)	3.96 – 5.52 (110)	1.39 – 2.60 (106)	82 – 152 (67)
Arithmetic mean (SD)	41 (8)	4.76 (0.25)	1.89 (0.23)	108 (14)
Geometric mean (Median)	40 (39)	4.76 (4.74)	1.88 (1.89)	107 (106)
Significance level	< 0.001	< 0.01	< 0.01	< 0.001

Figure captions

Fig. 1: Location of the DYFAMED sampling site (43° 25'N, 7°52' E), in the Ligurian Sea, with the North Current. The atmospheric time-series station (Cap Ferrat) is located close to Nice (see, e.g., Migon et al., 2008). Figure modified from Heimbürger et al. (2014).

Fig. 2: Temporal evolution of the mixed layer depth (MLD) and total integrated chlorophyll-a (ITChla) at the DYFAMED site from July 2007 to March 2009.

Fig. 3: Vertical (a) phosphate and (b) silicate concentration profiles (0-2200 m) at the DYFAMED site from July 2007 to March 2009.

Fig. 4: Vertical profiles of trace metals (0-2200 m) at the DYFAMED site from July 2007 to March 2009: a: Co, b: Ni, c: Cu, and d: Pb. Note Co and Pb concentrations are expressed in pM, and Ni and Cu concentrations are expressed in nM.

Figure 1

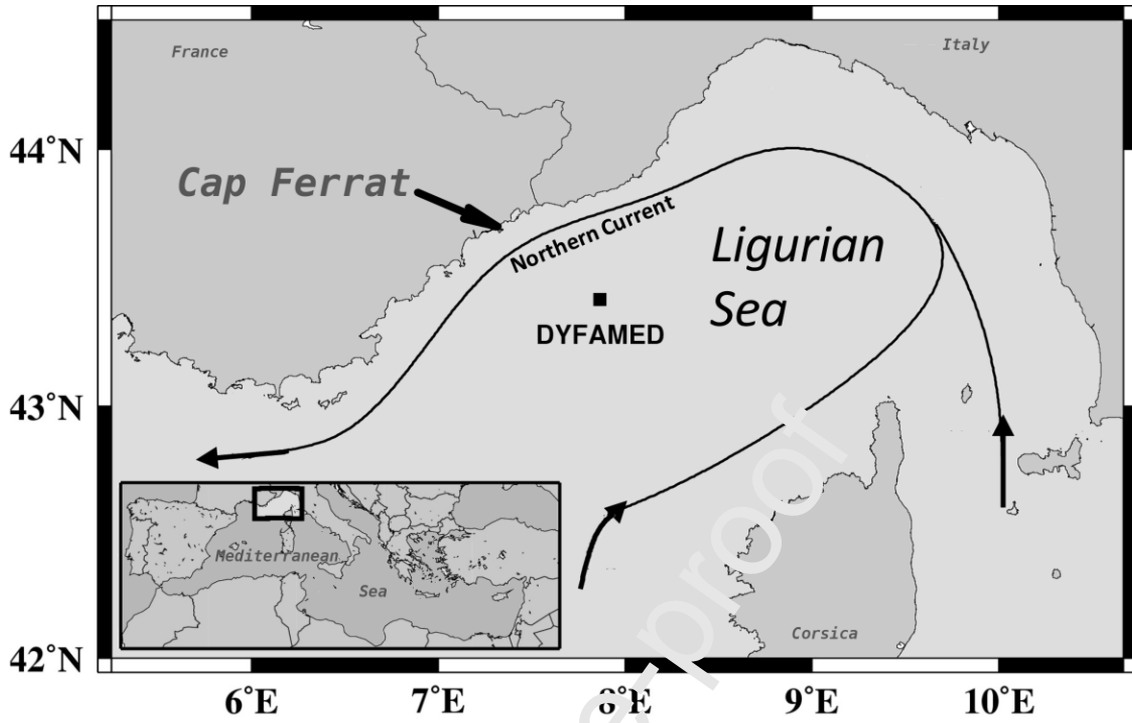


Figure 2

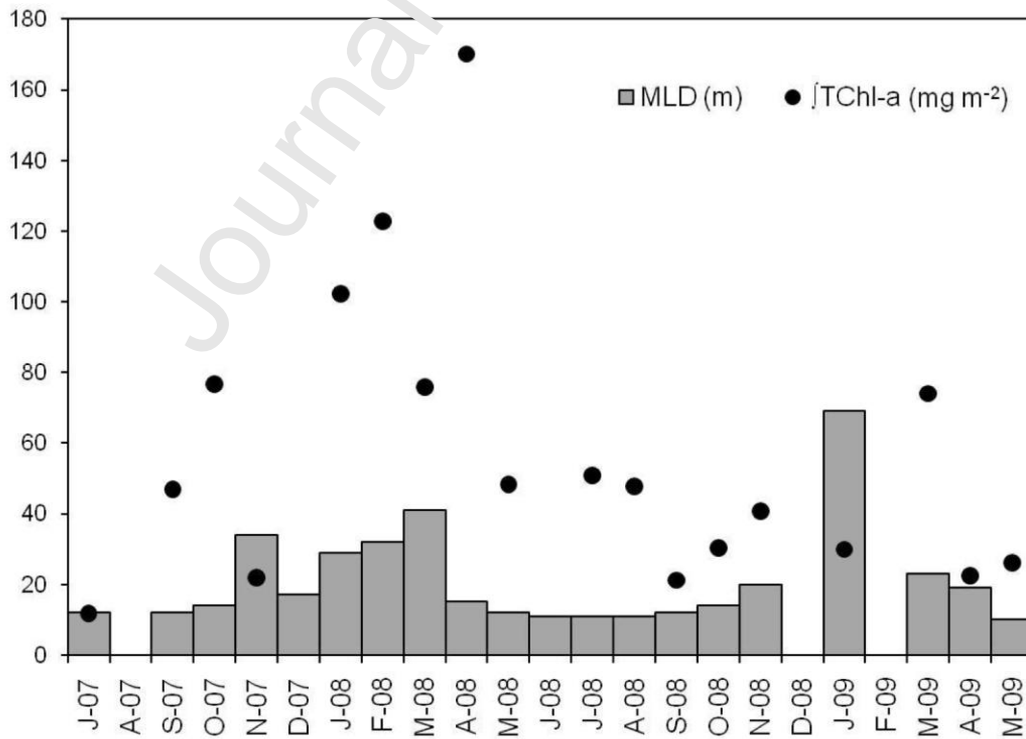


Figure 3

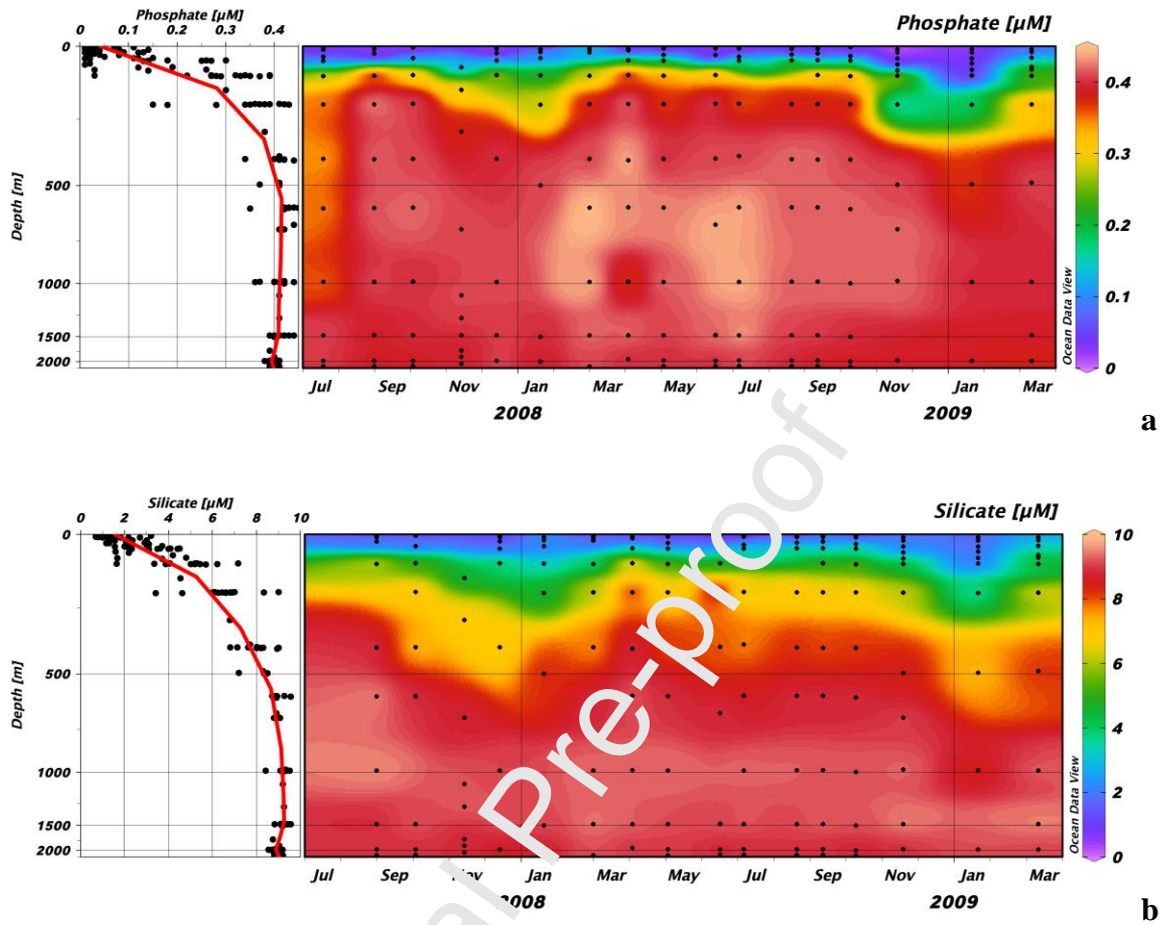
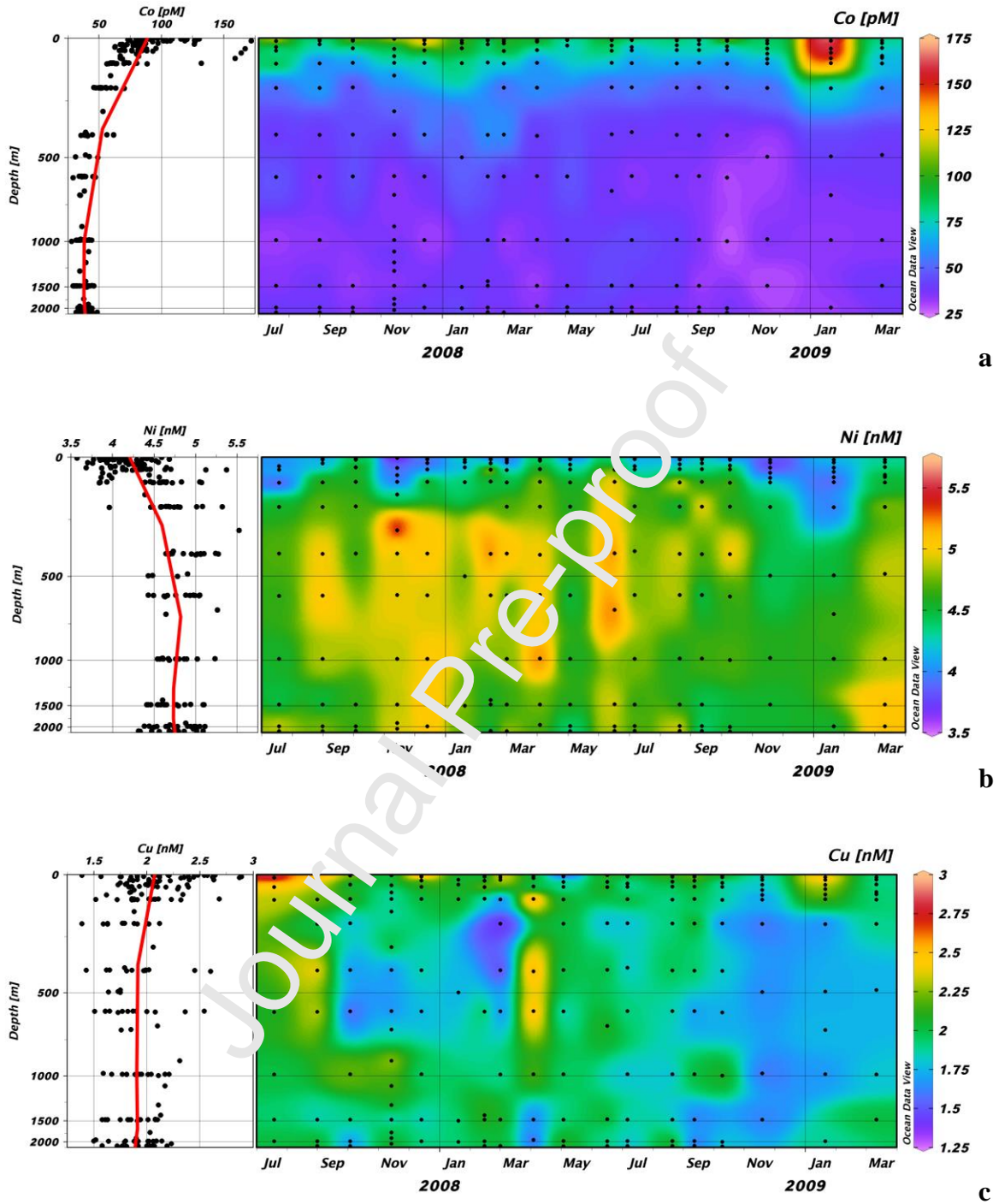
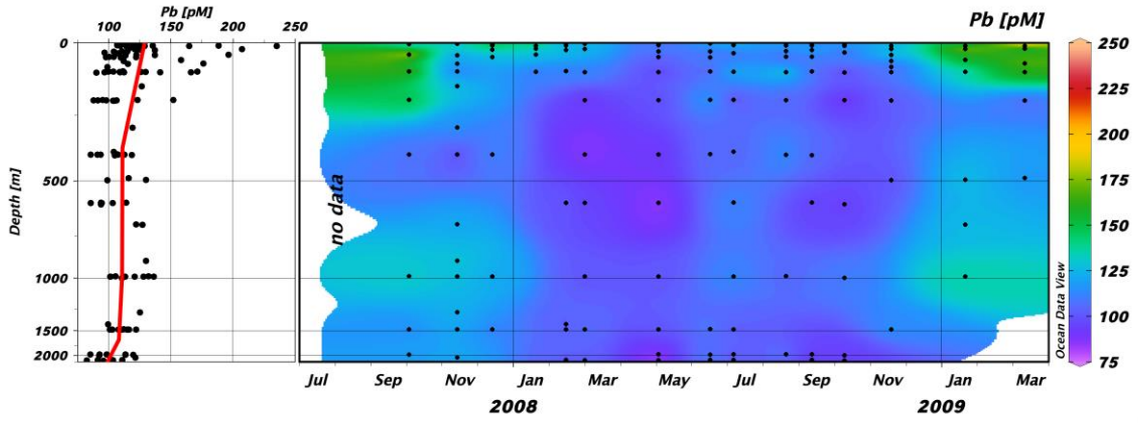


Figure 4





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Figure 1

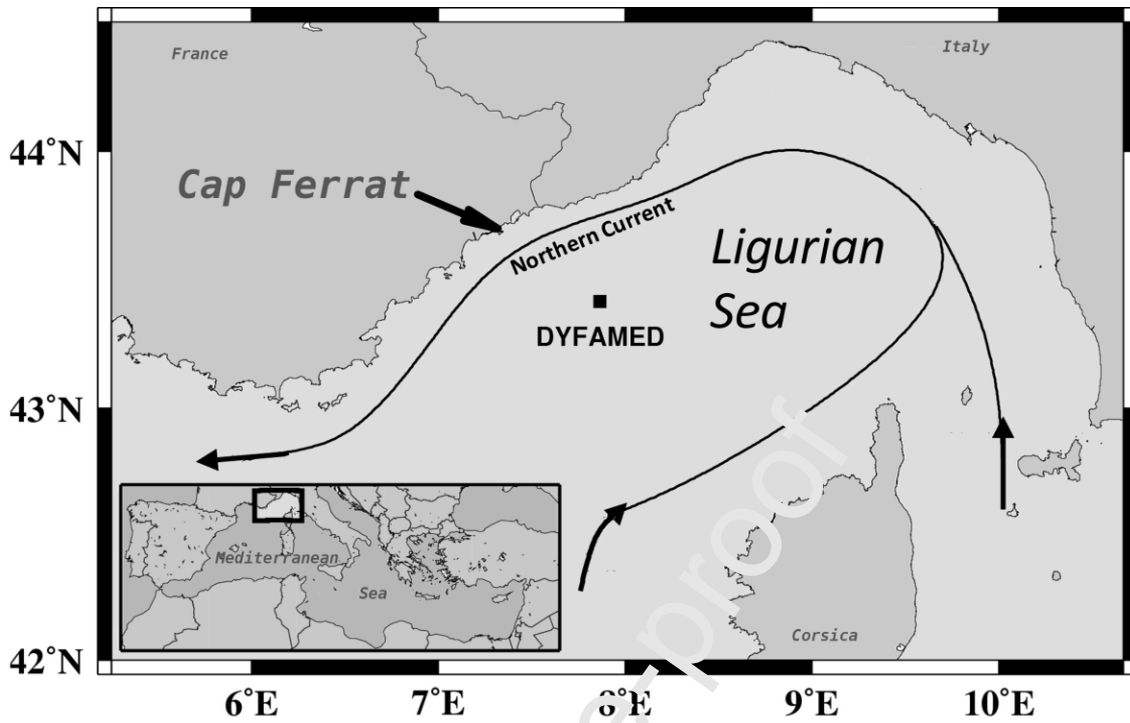


Figure 2

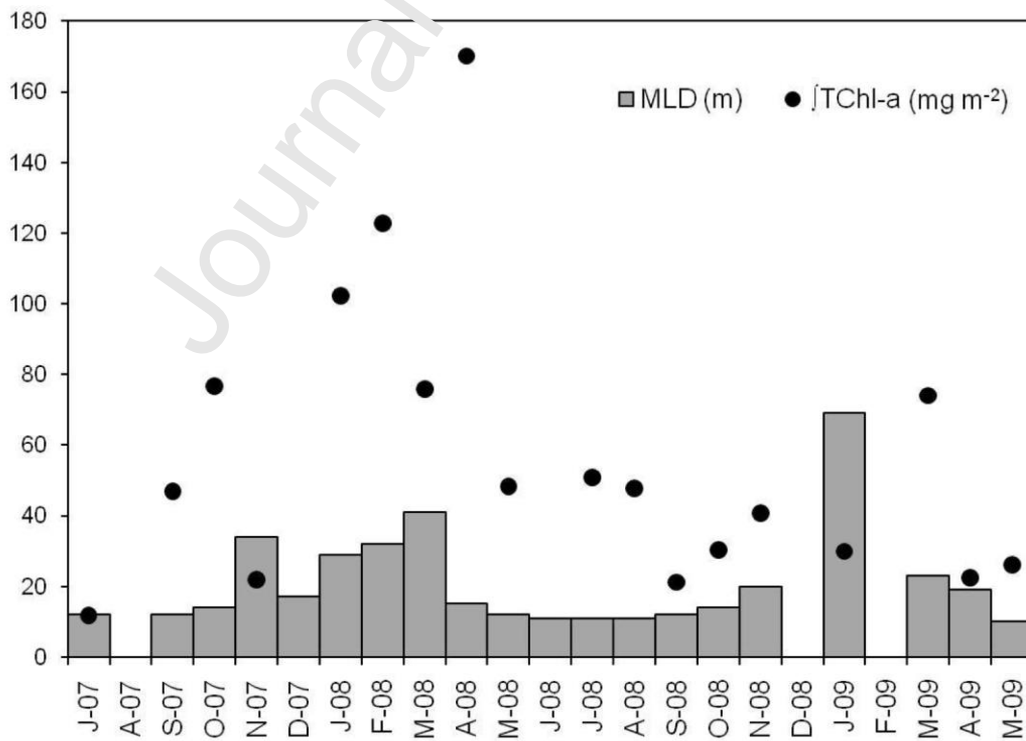


Figure 3

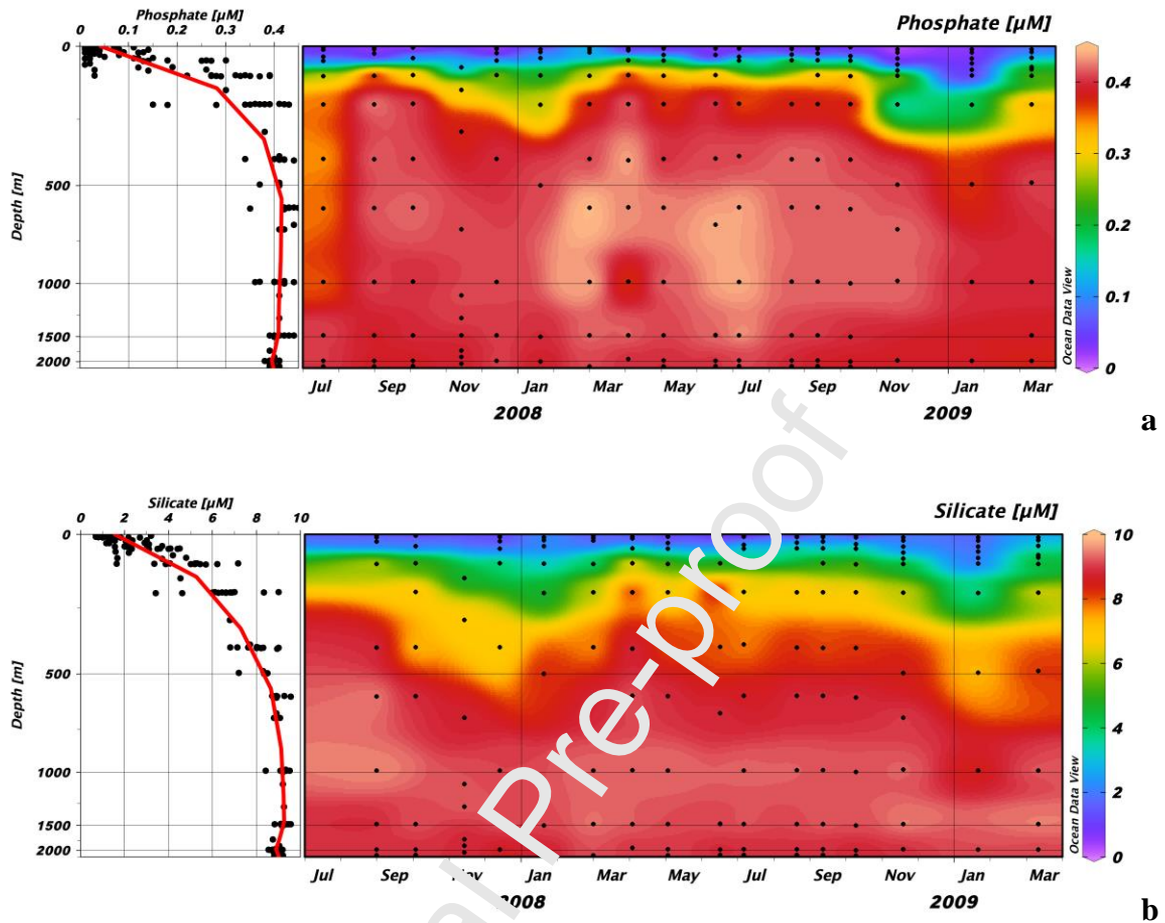
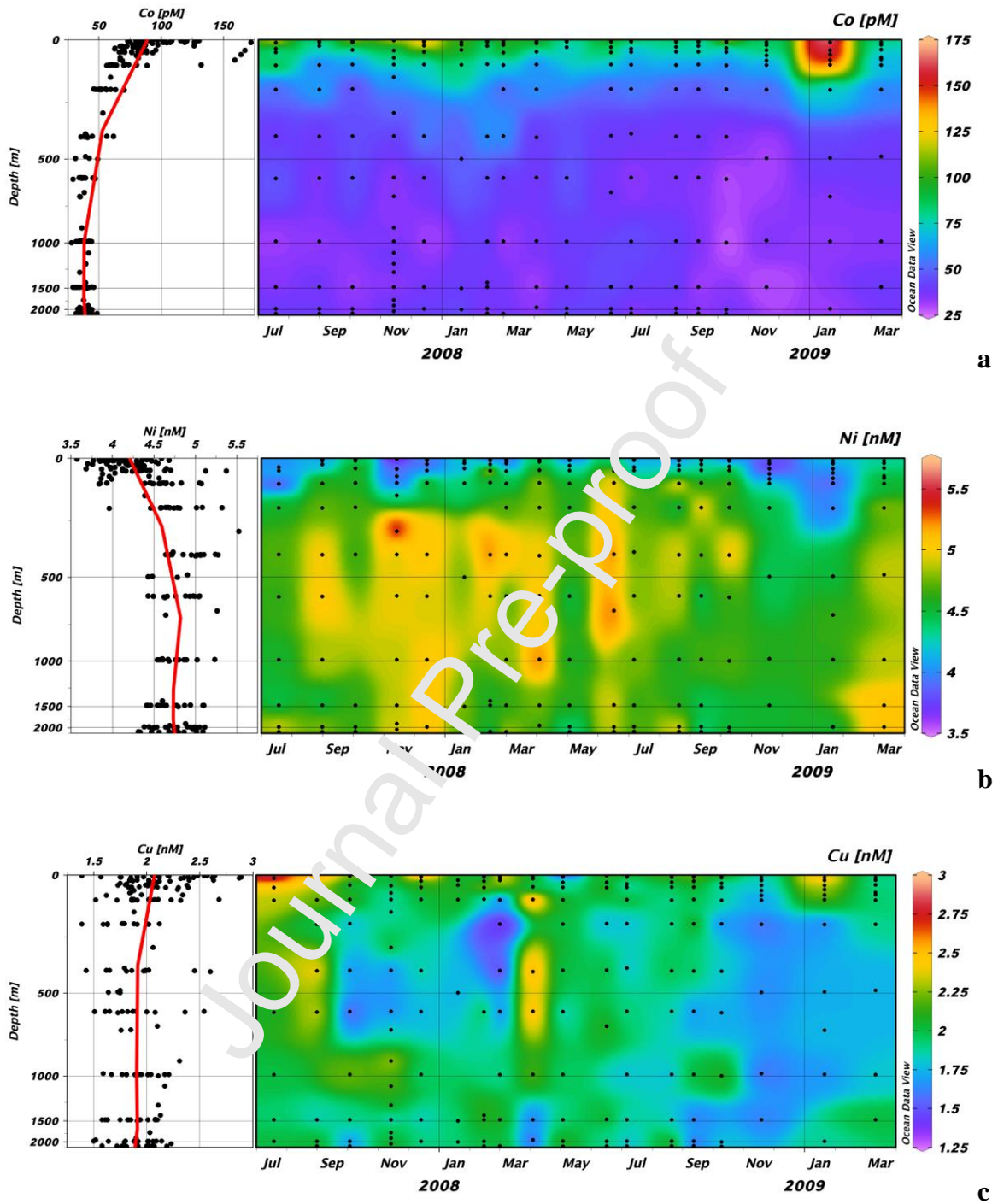
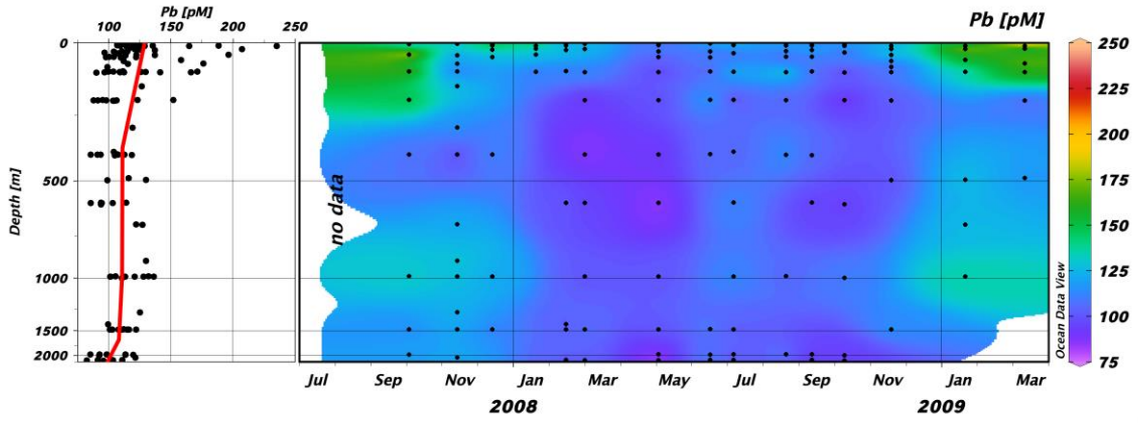


Figure 4





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Highlights

- We sort out the influence of these different parameters in the water column in a low-nutrient low-chlorophyll region, the Ligurian sea (Western Mediterranean);
- The intensity of winter mixing determines the homogenization of TM concentrations in the water column;
- TM exhibit a range of biogeochemical behaviours from surface-enriched (scavenged-type) to surface-depleted (nutrient-like) with Co and Ni as representative cases;
- Atmospheric input is a chief parameter as is determine Pb, Cu and Co distributions, and preclude any limitation of primary production by TM;
- Anthropogenic Pb decreased by up to 60% and 20% in surface and deep waters the last 20 years, but water column is still Pb contaminated, in such a way that the Western Mediterranean deep water outflow is the source of Pb contamination observed at 1000m in the adjacent Eastern North Atlantic Ocean.

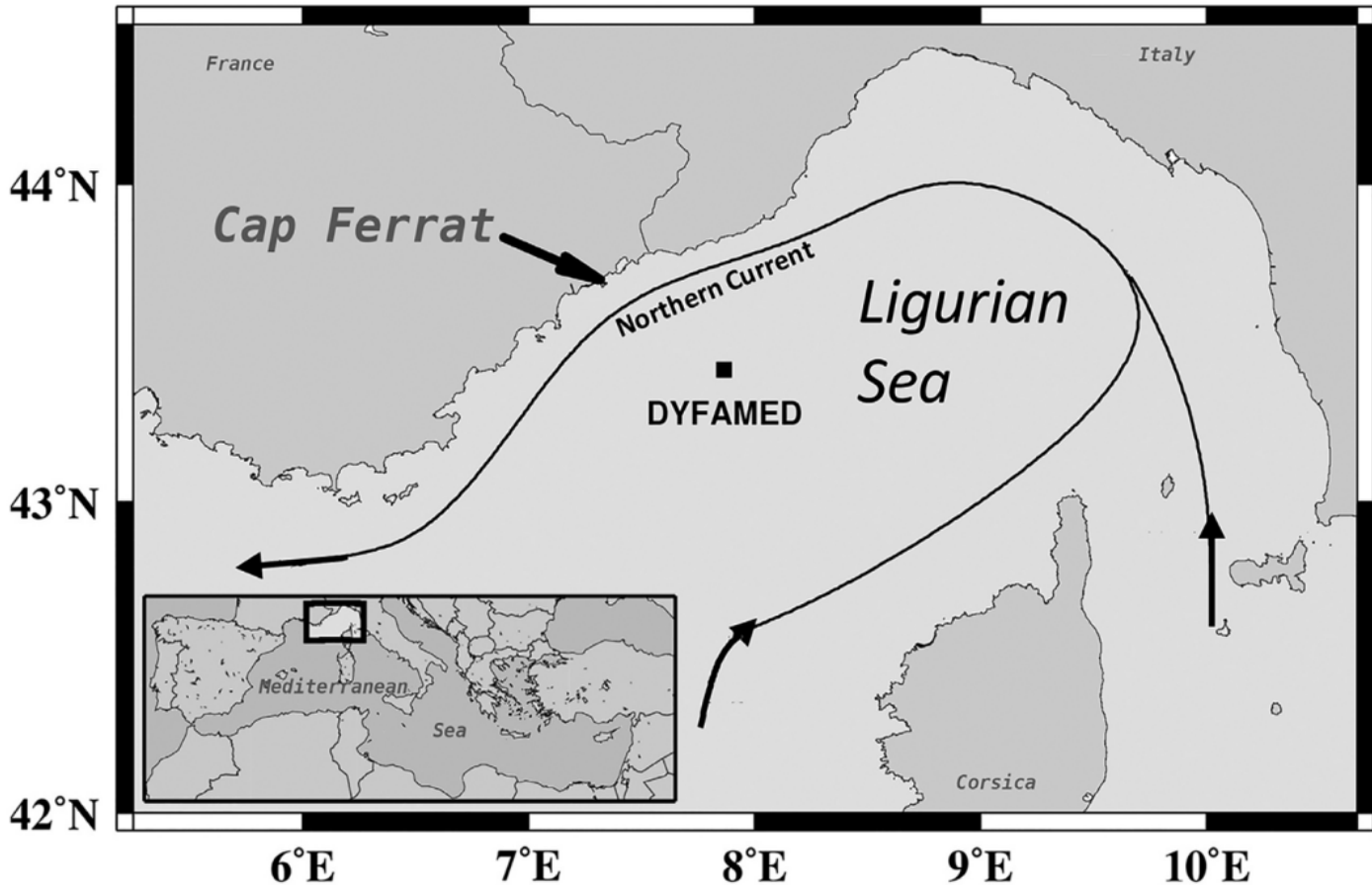


Figure 1

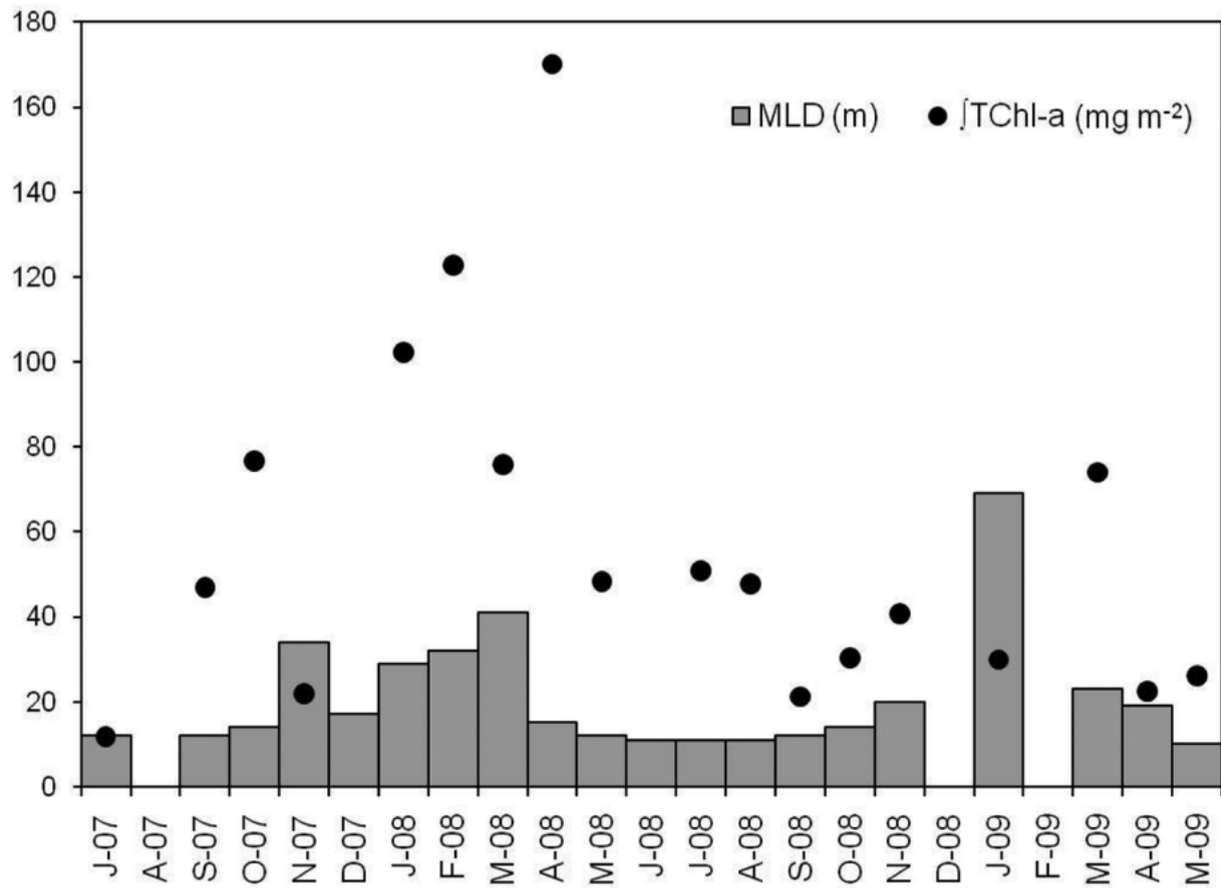


Figure 2

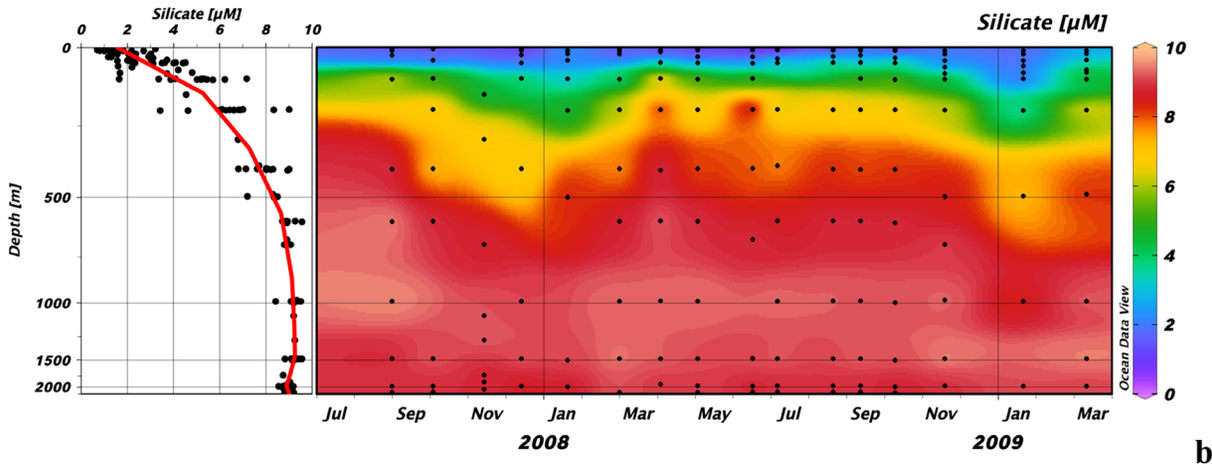
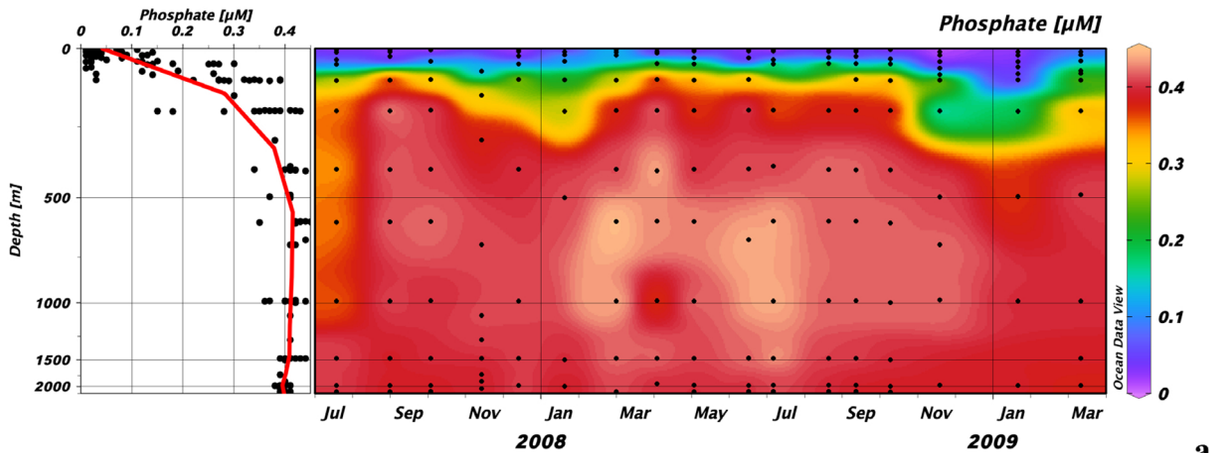


Figure 3

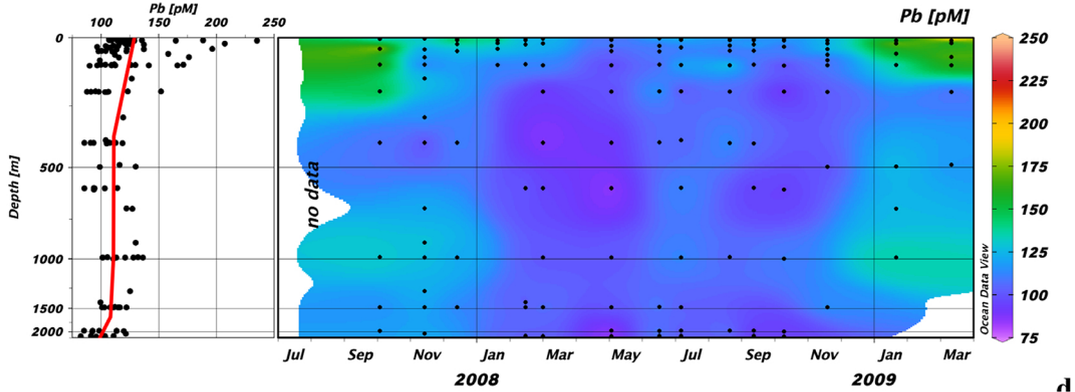
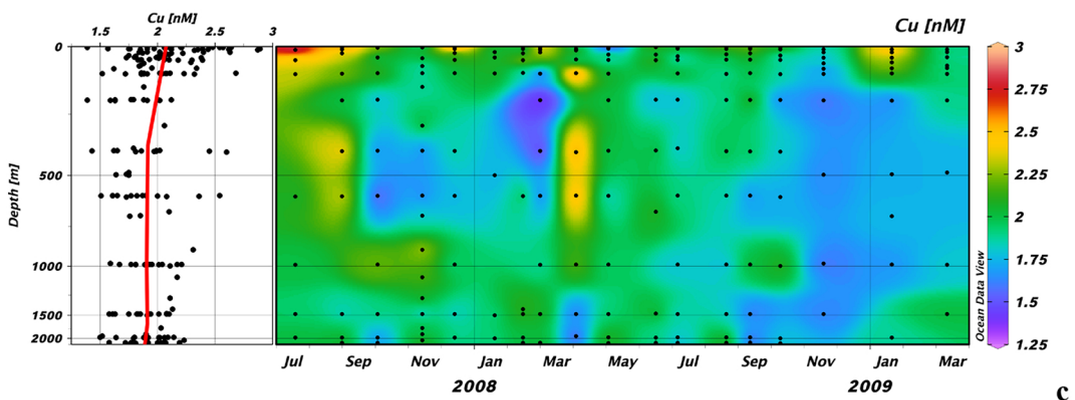
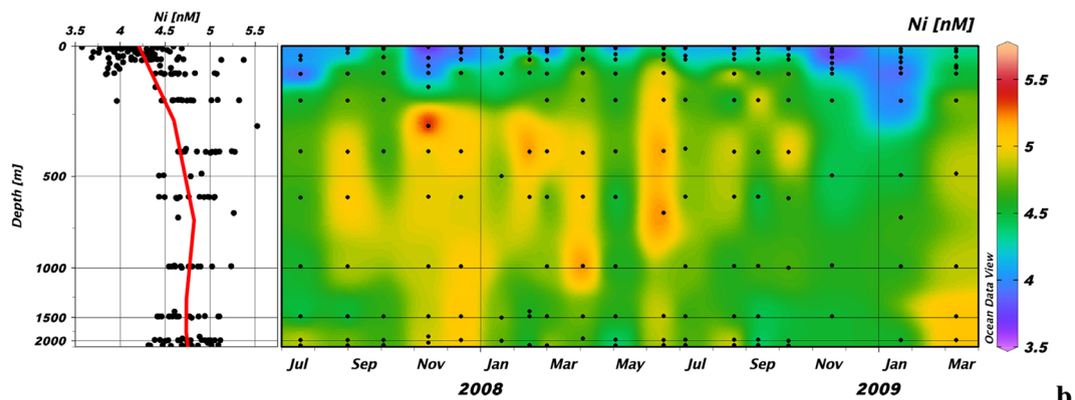
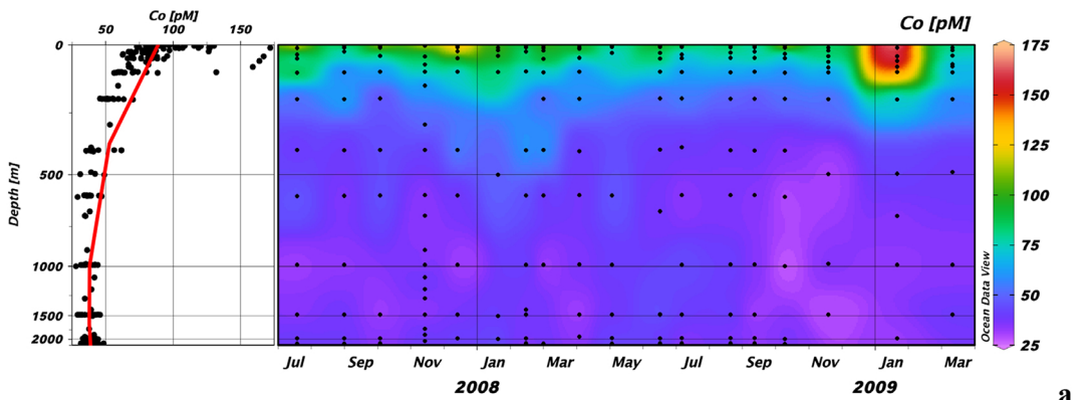


Figure 4